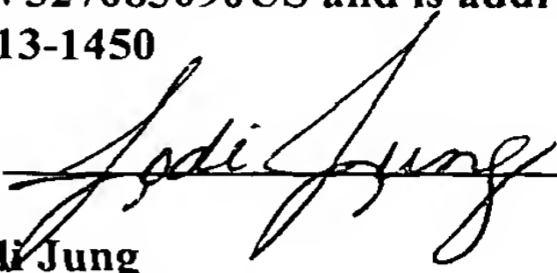

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SPECIFICATION

TO ALL WHOM IT MAY CONCERN:

25 BE IT KNOWN THAT WE, David W. Skinkle, a resident of Denver, Colorado, and a citizen of the United States of America; and Stephen W. Berch, a resident of Arvada, Colorado, and a citizen of the United States of America, have invented certain new and useful improvements in

30 **APPARATUS FOR SEPARATING BLOOD COMPONENTS**

of which the following is a specification.

APPARATUS FOR SEPARATING BLOOD COMPONENTS

Field of the Invention

This invention relates to an apparatus for separating components of whole blood. More particularly, this invention relates to an apparatus for the separation and collection of platelet poor plasma (PPP), platelet rich plasma (PRP), and red blood cells (RBC).

Background of the Invention

Whole blood can be collected from a donor and processed into different products. The collection and separation of blood typically has involved many steps as well as operator interaction.

Whole blood contains red blood cells, white blood cells, platelets, and plasma. Traditionally, these components were separated by a batch process in which a blood bag was spun for a period of approximately 10 minutes in a large refrigerated centrifuge. After centrifugation, the main blood constituents, red blood cells (erythrocytes), platelets and white blood cells (leukocytes) , and plasma sedimented and formed distinct layers. These constituents were then expressed sequentially by a manual extractor in different satellite bags attached to the primary bag.

More recently, automated extractors have been introduced. Nevertheless, the whole process remains laborious. There remains a widespread need for an apparatus that will automatically separate the different components of whole blood efficiently and easily.

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Summary of the Invention

The invention provides a centrifuge apparatus for processing blood comprising a bottom spring-loaded support plate; a top support plate; an axial inlet/outlet for blood to be processed and processed components of the blood, the axial inlet/outlet being attached to the top support plate by a rotating seal

assembly; a variable volume separation chamber mounted between the bottom support plate and the top support plate, the variable volume separation chamber being fluidly connected to the axial inlet/outlet; a pump fluidly connected to the axial inlet/outlet; and a rotary drive unit attached to the bottom support plate. The 5 top support plate is fixed vertically and the bottom spring-loaded support plate is mounted on springs that maintain pressure on the variable volume separation chamber and allow the bottom support plate to move vertically.

It is to be understood that both the foregoing general description and the following detailed description are exemplary and explanatory and are intended to 10 provide further explanation of the invention as claimed.

Brief Description of the Drawings

FIG. 1 is a perspective view of an apparatus of this invention.

FIGS. 2A and 2B are cross-sectional views of the centrifuge and FIG. 2C is 15 a cross-sectional view of an alternate centrifuge.

FIGS. 3A to 3C are cross sectional views of the rotating seal of the centrifuge.

FIGS. 4A to 4C are top, perspective and side views, respectively, of rigid support plate used in the centrifuge of FIGS. 2 and 3 and FIG. 4D is an exploded 20 view of the top support plate, port, and rotating seal assembly.

FIG. 5A is a perspective view and FIG. 5B is a side view of the cartridge that contains the fluid sensor and the valve assembly.

FIG. 6A is a side view and FIGS. 6B and 6C are cross-sectional views of the fluid sensor pathway.

25 FIGS. 7A and 7B are cross-sectional and side views, respectively, of the valve assembly when whole blood is flowing into the centrifuge.

FIGS. 8A and 8B are cross-sectional and side views, respectively, of the valve assembly when PPP is flowing out of the centrifuge.

30 FIGS. 9A and 9B are cross-sectional and side views, respectively, of the valve assembly when platelet rich plasma is flowing out of the centrifuge.

FIGS. 10A and 10B are cross-sectional and side views, respectively, of the valve assembly when RBC and air are flowing out of the centrifuge and into the RBC compartment of the reservoir.

5 FIGS. 11A and 11B are cross-sectional and side views, respectively, of the valve core.

FIG. 12 is a cut-away partial detail view of the housing of the apparatus with the 4-way valve cartridge attached.

FIG. 13 is a partial cross-sectional view of the cartridge snapped on the housing.

10 FIG. 14 is a side view of the multicomponent 3-compartment reservoir bag.

FIG. 15 is a perspective view of an alternate embodiment of the apparatus of this invention.

FIGS. 16A and 16B are perspective views of a pinch valve.

15 FIG. 17 is a perspective view of the disposable component of the apparatus of FIG. 15.

Description of the Preferred Embodiments

In one embodiment, the invention provides an apparatus comprising a centrifuge with spring-loaded plate and top locking feature, a valve driver mechanism, fluid sensor, peristaltic pump and touch screen computer interface. Additionally, a syringe chiller may be provided to keep various components in a syringe at a desired temperature.

In a preferred embodiment, a single use, sterile disposable processing set interfaces with the apparatus. The sterile disposable consists of a circular variable volume separation chamber with axial rotating seal assembly, 4-way valve cartridge with integral sensor and fluid pump loop and a pre-attached three compartment reservoir bag. The three-compartment reservoir bag consists of a chamber for anticoagulated whole blood, a chamber for platelet poor plasma, and a chamber for concentrated red blood cells. Platelet rich plasma is collected in a sterile syringe attached to the 4-way valve luer lock port.

Specifically, the invention provides a centrifuge apparatus for processing anticoagulated whole blood comprising a bottom spring-loaded support plate, a slotted top locking feature, and a stator arm assembly. The disposable variable volume separation chamber with rigid support plate loads and locks into the spring loaded centrifuge chamber. The rotating seal of the separation chamber is interfaced and held stationary by the stator arm assembly. Tubing is attached to the rotating seal assembly to provide an axial inlet/outlet for blood to be processed and processed components of the blood. The 4-way valve assembly is attached to the inlet/outlet tube of the variable volume separation chamber. The 4-way valve is mounted to the fluid sensor and snaps to the top housing of the apparatus.

5 Rotation of the peristaltic pump loads the fluid pump loop. The three compartment bag is attached to side of the apparatus to allow access to fluid inlet and outlet ports.

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In a preferred embodiment, this invention achieves separation of whole blood components according to the following method of operation. The disposable processing set is attached to the apparatus. Whole blood collected from the donor is mixed with anticoagulant and delivered to the inlet port of the reservoir bag whole blood compartment. The clinician selects the desired whole blood volume to process on the user interface. The start button is selected to initiate the separation cycle and rotation of the centrifuge. The valve driver positions the 4-way valve to the whole blood compartment and the peristaltic pump drives fluid from the reservoir to the spinning variable volume separation chamber. Fluid pressure inside the rotating separation chamber increases with increased gravitational force and the addition of whole blood. This pressure drives the spring load bottom plate downward allowing additional volume to enter the rotating system. The flexible variable volume separation chamber changes shape and this shape change is limited by a fixed stop internal to the centrifuge housing. Once adequate separation of the whole blood components occurs, the centrifuge rotation speed is decreased. The peristaltic pump direction is reversed, pumping the component layers from the axial port of the separation chamber. The apparatus

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fluid sensor detects the concentration of the various component layers and utilizes algorithms to change the 4-way valve position to the desired component layer collection vessel. The process is complete when all component layers are collected and the apparatus fluid sensor senses air. The 4-way valve fluid path
5 allows the draw back of platelet poor plasma from the PPP reservoir compartment into the PRP syringe. Multiple whole blood separation cycles are possible with this invention.

The advantages of this invention include the use of an automated system and the ability to separate variable quantities of blood. Even very small quantities
10 of whole blood can be efficiently separated, collected, and returned to a patient using the apparatus of this invention. Larger volumes can also be selected and processed within approximately the same cycle time of smaller volumes, allowing the clinician to harvest a larger quantity of platelets per cycle. This is advantageous in patients with low platelet counts where more whole blood can be
15 collected and processed in approximately the same cycle time with less dilution of the PRP product to produce substantially higher baseline multiples.

Additional advantages include the use of the fluid sensor to produce a PRP and PPP product void of red blood cells. The first component layer removed from this apparatus after centrifugation is the PPP layer. The fluid sensor detects when
20 the PPP product is clear and free of red cells prior to collection. The same is true for the PRP collection. Once platelets are sensed the platelet collection process is initiated and continues until red blood cells are sensed. The user can predetermine the concentration of red blood cells in the final PRP product. This is advantageous in certain clinical procedures.

25 The invention provides a centrifuge apparatus for processing blood comprising a bottom spring-loaded support plate; a top support plate; an axial inlet/outlet for blood to be processed and processed components of the blood, the axial inlet/outlet being attached to the top support plate by a rotating seal assembly; a variable volume separation chamber mounted between the bottom
30 support plate and the top support plate, the variable volume separation chamber

being fluidly connected to the axial inlet/outlet; a pump fluidly connected to the axial inlet/outlet; and a rotary drive unit attached to the bottom support plate. The top support plate is fixed vertically and the bottom spring-loaded support plate is mounted on springs that maintain pressure on the variable volume separation chamber and allow the bottom support plate to move vertically.

The invention provides a method of processing blood comprising: providing a centrifuge apparatus as described above; introducing a quantity of blood into the variable volume separation chamber; centrifuging the blood; and removing the separated components of the blood through the axial inlet/outlet.

The invention also provides a centrifuge apparatus for processing blood comprising a bottom support plate; a top support plate; an axial inlet/outlet for blood to be processed and processed components of the blood, the axial inlet/outlet being attached to the top support plate by a rotating seal assembly; a variable volume separation chamber mounted between the bottom support plate and the top support plate, the variable volume separation chamber being fluidly connected to the axial inlet/outlet; a pump fluidly connected to the axial inlet/outlet; and a rotary drive unit attached to the bottom support plate. The top holder is fixed vertically and the bottom support plate is mounted on a ball-screw actuator that maintains pressure on the variable volume separation chamber and allows the bottom support plate to move vertically.

The invention provides a disposable cartridge comprising a plurality of ports for receiving or dispensing blood or blood components and a fluid sensor pathway for displaying blood or blood components for analysis, the cartridge being adapted to be mounted on a multi-position valve for directing flow between the ports and the fluid sensor pathway being adapted to be mounted adjacent to one or more sensors for analyzing blood.

The invention provides a disposable set comprising: a container for blood; a plurality of containers for receiving separated components of the blood; a disk-shaped bag; a top support plate for a centrifuge; an axial inlet/outlet for blood to be processed and processed components of the blood, the axial inlet/outlet being

attached to the top support plate by a rotating seal assembly; and tubing. The disposable set may further comprise a disposable cartridge comprising a plurality of ports for receiving or dispensing blood or blood components and a fluid sensor pathway for displaying blood or blood components for analysis, the cartridge being
5 adapted to be mounted on a multi-position valve for directing flow between the ports and the fluid sensor pathway being adapted to be mounted adjacent to one or more sensors for analyzing blood.

The blood component separation apparatus, as shown and described in the Figures, includes housing 10 containing centrifuge 20 (shown in cross section in
10 FIGS. 2A and 2B). With reference to FIG. 1, the apparatus includes a user interface that comprises touch screen display 30. A single power supply is used for the centrifuge motor and for the electronics. The apparatus also includes a fluid sensor 60, 4-way valve 70, and 3-compartment reservoir/collection bag 90. The fluid sensor pathway 60 and the 4-way valve 70 are contained within cartridge
15 80, which in a preferred embodiment has a snap-on feature so that it is readily removable and maintains its position on the apparatus during use. Blood and its components flow to and from the 3-compartment reservoir/collection bag by means of flexible tubing. A syringe chiller 85 may be provided to keep various fluids in a syringe at a desired temperature. For example, a Peltier device can be
20 used to cool or heat the syringe.

Blood is withdrawn from a patient, mixed with an appropriate anticoagulant (ACD-A, CPD-A) and placed in compartment 906 of the 3-compartment reservoir/collection bag 90, as illustrated in FIG. 1. The bag is connected via tubing line 416 to the 4-way valve core 71 contained within the disposable
25 cartridge 80 (FIGS. 11A and 11B). The disposable cartridge includes a tubing loop 411, which is configured to fit within the raceway of the peristaltic pump 40 contained on the top surface of the housing. One end of the tubing connects to an inlet of a flexible and variable volume separation chamber. The blood is pumped into the centrifuge, where, in the flexible separation chamber, it is separated into
30 concentrated red blood cells (RBC), platelet poor plasma (PPP) and platelet rich

plasma (PRP). These components are then transferred out of the flexible separation chamber via tubing lines 410 and 411. The RBC component is transferred to compartment 904 via tubing line 414, and PPP flows into compartment 902 via tubing line 412. The platelet rich plasma (PRP) flows into a 5 syringe via port 802 (syringe not seen in FIG. 1).

The flexible variable volume separation chamber 140 and top support plate 114 fits within the spring loaded plate 110 and top 111, having locking feature 112, on the centrifuge assembly 20. A stator arm assembly 113 engages rotating seal 120. A spring-loaded support plate 110 presses upward against the variable 10 volume separation chamber 140 though it is to be understood that the chamber and holder could be configured so that movement of the plate could be in any desired direction. Motion of the plate, rotation of the peristaltic pump, specified whole blood volume and reduced rotational speed causes expulsion of blood components. These components can exit the port 124 coincident with the axis of the rotating 15 seal assembly 120. A lid 115 covers the centrifuge.

The valve system coupled with optical sensors permits the automation of this process. The graphical user interface (GUI) is object oriented and uses a unified modeling language. The apparatus thus can be used by operators who have varying levels of sophistication.

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Centrifuge

In operation, whole blood from the 3-compartment reservoir 90, specifically compartment 906 is pumped through the valve into a variable volume separation chamber 140. The centrifuge is then rotated to separate the blood components. 25 The heavier components migrate to the outer portions of the separation chamber while the lighter components remain near the center of the separation chamber . Centrifuge 20 is shown in cross section in FIGS. 2A and 2B.

Motor 102 is operably connected to hollow shaft 104 which is integrally formed with or mounted on spring-loaded support plate 110. Coil springs 106 30 comprise one or more springs fit onto shafts 107 and are operably connected to

support plate 110. Rotating seal assembly 120 includes port 124. In FIG. 3B, lip-seals 301 attached to center hub 300 seal against a stationary disk 302 which forms a fluid seal. Port 124 provides a passageway to the variable volume separation chamber 140, which is held in the space between rigid support plate 114 and spring loaded plate 110. Motor 102 rotates spring loaded plate 110, rigid support plate 114, and variable volume separation chamber 140.

During centrifugation, lower density blood components accumulate in the center region 145 of separation chamber 140, that is, close to the axis of rotation, while higher density components are urged toward the outermost region.. The bottom support plate 110 moves down to accommodate the blood components due to centrifugal force.

For example, once whole blood has filled the separation chamber in the centrifuge, the centrifuge is run for 7 minutes at 4000 rpm. Then the rotation of the centrifuge motor is decreased. Decreasing the speed of the centrifuge causes reduced pressure inside the bag, allowing the spring-loaded support plate 110 to move upward against flexible reservoir 140, causing its contents to be expelled via port 124. This, along with operation of peristaltic pump 40 in a direction reverse to that during which the variable volume separation chamber was filled, causes expulsion of the blood components through the fluid exit port coincident with the axis of the centrifuge (i.e., port 124).

Because PPP is less dense, it is expelled first. The PPP is directed through tubing 410 to the valve system, and fed into the PPP compartment of the 3-chamber reservoir bag., as described below for FIGS. 7 to 10. Other components follow in sequence, also as described further below.

FIG. 2C illustrates an alternate embodiment of the centrifuge, and shows centrifuge 20b in a cross-sectional view wherein ball-screw actuator 206 is used to control the space between the top holder and the bottom plate. The ball-screw actuator 206 as an example is automatically controlled to maintain the desired space between the top holder 114 and the bottom plate 110. It is possible to use other mechanical means such as pneumatic, hydraulic or other mechanical

actuators to achieve desired positioning of the bottom plate. Use of a pump to move fluids could be eliminated with a mechanical actuator. Such an actuated bottom plate could draw fluids in and expel fluids from the separation chamber.

Typically the apparatus of this alternate embodiment invention will be used
5 by placing whole blood in the 3-chamber reservoir bag and transferring it from there to the centrifuge. However, it is possible to collect blood directly from the patient in the flexible disk bag of the centrifuge. The flexible disk bag is the preferred embodiment of the variable volume separation chamber. Since it is necessary to mix the collected blood with an anti-coagulant, it is important to
10 know the amount of blood being processed. The flexible disk bag is mounted in the centrifuge between the top plate and the bottom plate. The distance between the top plate and the bottom plate correlates to a known volume of blood in the flexible disk bag as shown by a graduated scale 117 (shown in FIG. 18), which the user can visually inspect to determine the amount of blood entering the flexible
15 reservoir. The operator thus can monitor the amount of anti-coagulant and whole blood collected without the use of a separate scale.

FIGS. 3A to 3C show various embodiments of a rotating seal. The rotating seal shown in FIG. 3A is equivalent to the one shown in FIG. 3B. The only difference in the designs is the presence of two bearings 303 in FIG. 3A versus one bearing in FIG. 3B. Two bearings provide greater stability of rotating components.
20 FIGS. 3A and 3B both have a center hub 300 that is attached to the center axis of the separation chamber 140. Bearing 303 is press fit onto the shaft of the center hub 300. Flexible lip-seals 301 are attached to the center hub 300 and create a fluid seal with a stationary sealing disk 302. Disk 302 is centered and sealed by a
25 compressible gasket 304 that also provides a pre-load to the outer race of the bearing 303 when compressed. The lower housing 305 provides features that contain the above mentioned components. Top housing 306 is attached to the lower housing 305 and compresses and seals the internal components of the rotating seal. Prior to assembly the center tube 307 is attached to the upper
30 housing either by press fit, insert molding or molded. The center tube allows for

the transfer of fluid in and out of the device. The rotating components of this design are the separation chamber 140, the center hub 300, and the lip seals 301. All other components are held stationary during centrifugation. The rotating lip-seal 301 interface to stationary disk 302 creates a positive and negative pressure

5 fluid seal.

FIG. 3C is similar to FIGS. 3A and 3B. The design shown in FIG. 3C differs with a press fit bearing 303 into the lower housing 305, a one piece lip seal 310 versus two lip seals and a flexible gasket 311 that seals the outer edge of the sealing disk 302. The upper housing 306 is attached to the lower housing 305 and

10 compresses gasket 311.

FIGS. 4A to 4C show various views of the top support plate 114. The top support plate is designed to provide structural support for the expanding variable volume separation chamber. FIG. 4D illustrates an exploded view of the separation chamber assembly, showing the top support plate 114, port 124, rotating seal assembly 120, center hub 300 and variable volume separation chamber 140. The rotating union generated by the rotating seal assembly 120 allows fluid to enter and exit the variable volume separation chamber while the chamber is rotating.

15 20 Graphical User Interface

Housing 10 includes a user interface comprising touch screen display 30, a stop button 301, a power switch, and various connectors for external electrical interface. The touch screen is resistive so that it will function if the operator is wearing gloves. The stop button is used to interfere with automatic operation if the operator deems necessary. All other operator interfacing is accomplished from this one screen using 3-D appearance of control features and judicious use of color. The external interfaces are used to upgrade software, download data, and possibly connect to a printer.

Cartridge

FIG. 5A shows cartridge 80 that contains fluid sensor pathway 60 and the valve assembly 70. Cartridge 80 typically comprises injection molded polymer. The cartridge is provided with snap-on features 82 and 84 that permit easy removal
5 of the cartridge from the apparatus. (Snap tab 890 is shown in greater detail in FIG. 13.) Valve housing 870 contains ports 802, 804, 806, 808, and 810, each of which connect to tubing lines for the transfer of fluid to and from the centrifuge to the 3-compartment reservoir and/or to a syringe. Cartridge 80 also includes fluid sensor pathway 60. Tubing 62 connects port 810 to fluid sensor pathway outlet
10 61a. Tubing 62 is placed in the raceway of the peristaltic pump. Blood flows in and out of fluid sensor pathway 60 through outlets 61a and 61b.

FIGS. 6A to 6C illustrate the section views of the fluid pathway geometry for the fluid sensor. FIG. 6B provides a side section view of the fluid sensor pathway in which the fluid flow is thinned out to allow improved detection of
15 whole blood component layers.

Sensors

Blood components flow through the fluid sensor pathway 60 and the flow is monitored at various wavelengths. An algorithm is used to determine what
20 component layer is in the fluid sensor pathway: RBC, PPP, PRP, or air. A combination of absorption and scattering causes the signal to change. Computer software controls the intensity of the LEDs. A cutaway view of the photodiode detector placement is shown in FIG. 12. Photodiode detectors are configured to fit in pockets or openings 510 and 512. These pockets are adjacent to fluid sensor
25 pathway 60.

The system comprises three LEDs and two photodiode detectors. The first LED is infrared, emitting light with a wavelength of 1300 nm. This LED is matched to one of the two photodiodes. The second LED is also infrared, emitting light with a wavelength of 940 nm, and the third LED is blue, emitting light with a
30 wavelength of 470 nm. The second photodiode is used to detect the light energy

from the second and third LEDs. The second photodiode is more responsive to the 940 nm light. Accordingly, the 470 nm LED is set so that it shined directly at the detector, while the 940 nm LED is positioned off center.

The blood path through the sensor is a rigid polycarbonate piece with a near elliptic cross section. The light shining through the piece sees the blood interface on a flat surface. The light is directed normal to that surface. The light emitted from each LED is electronically chopped by pulsing the LEDs on and off in sequence. The detector response is then sampled so that any signal due to the ambient background light can be cancelled out.

The intensity of the light emitted from the LEDs is electronically adjustable through a current sensing, voltage feedback amplifier. The signal from the detector is monitored, while the intensity of the light is adjusted, until the signal falls within a pre-defined window. This process is accomplished automatically in software for every new case performed on the machine. The operator is not involved in any way with this calibration process.

The different blood components are identified by considering the intensity of the light transmitted through the blood, as well as the derivative of the intensity as a function of time. Because the blood is flowing through the sensor while the light intensity is being sampled, the derivative of the intensity is also a function of the blood volume passing through the sensor. The components that are identified are: (1) whole blood, (2) platelet poor plasma, (3) platelet rich plasma, (4) red blood cells, and (5) air .

Valve

The blood is separated into components in the centrifuge, which is connected by tubing to a 4-way valve 70 contained within a disposable cartridge 80 (FIGS. 5 to 11). The disposable cartridge can be snapped onto the housing for ease of removal.

The valve is designed so that whole blood flows through the valve in its “home” position and into the centrifuge, as illustrated in FIGS. 7A and 7B. The

valve is then rotated to the position shown in FIG. 10A to allow the pump to pump air through the tubing to displace the residual whole blood volume into the centrifuge. After centrifugation of the whole blood, the valve moves to provide a path for the platelet poor plasma (PPP) to exit the centrifuge (FIGS. 8A and 8B).

5 The valve then moves so that platelet rich plasma (PRP) can flow from the centrifuge into the collection bag (FIGS. 9A and 9B). In FIGS. 10A and 10B, red blood cells (RBC) exit the centrifuge. Also in this position while red blood cells are being pumped from the separation chamber, platelet poor plasma (PPP) can be aspirated back to be mixed with platelet rich plasma (PRP) to dilute the PRP, if
10 desired.

FIGS. 11A and 11B show the 4-way valve core 70. A stepper motor drives the valve core 70 to the desired locations during processing. There is a potentiometer to verify valve position based on voltage. Voltages and current are measured for diagnostic purposes.

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Reservoir/collection bag

FIG. 14 shows an embodiment of 3-compartment reservoir bag 90. The bag is flexible and preferably comprises a PVC polymer. Preferably the bag is transparent so that fluid levels can be seen easily. Whole blood is placed into compartment 906. The blood is pumped out of this compartment (by peristaltic pump 40) and into the variable volume separation chamber 140 of the centrifuge. Reservoir/collection bag 90 is shown as a 3-compartment unit, and it is to be understood that this bag could be perforated for easy separation of the compartments.

25 After centrifugation, the blood components are sent from the centrifuge through tubing line 410 through peristaltic pump 40 via tubing line 411 into valve 70 and thus to the separation/collection bag via tubing lines 412, 414, and 416. See FIG. 1. A preferred embodiment of the bag is shown in FIG. 14, wherein PPP is in compartment 902, RBC is in compartment 904, and whole blood is in
30 compartment 906. Compartments 902 and 904 are provided with outlets 912 and

914 so that the contents can be completely and easily removed, typically by syringe. Compartment 906 is provided with inlet 913. In addition, compartments 902, 904 and 906 are provided with vents 924 and 926 to allow air to enter and exit.

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Alternate embodiments

An alternate embodiment of this invention is illustrated in FIGS. 15 to 17. This apparatus functions similarly to the apparatus described above in FIGS. 1 to 14. FIG. 15 shows housing 1000 with display 1030. A separate reservoir 1096 for whole blood is provided and connected via tubing line 1046 to a common tubing line 1048 that is operably connected via pump loop 1040 to a pump (not shown) and to the centrifuge (not shown). Whole blood is sent to a variable volume separation chamber in the centrifuge where it is separated (as described above for the previous embodiment) and then the PPP and RBC components are sent to the reservoirs 1092 and 1094, respectively. PRP is collected by means of syringe 1098.

Fluid sensor pathway 1060 and sensors 1050 are directly in the line to/from the centrifuge. Three pinch valves 1072, 1074, and 1076 are provided in lines 1042, 1044, and 1046 that lead from the PPP, RBC, and WB reservoirs, respectively. These valves are operably connected to the sensors thereby automatically sending the desired fluid to the correct reservoir. For example, pinch valve 1072 is shown in FIGS. 16A and 16B. Pinch valve 1702 comprises solenoid 1700, pinch spring 1703, occluding edge of valve 1704, position sensor 1706, and housing 1710. FIG. 17 illustrates a disposable portion of apparatus 1000.

The above description and the drawings are provided for the purpose of describing embodiments of the invention and are not intended to limit the scope of the invention in any way. It will be apparent to those skilled in the art that various modifications and variations can be made without departing from the spirit or

scope of the invention. Thus, it is intended that the present invention cover the modifications and variations of this invention provided they come within the scope of the appended claims and their equivalents.